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(A) Destructurized starch and procees for making eams.

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 and that its mass everage moler mass is reduced by a factor of 2
 5000 compared to 10 original mass average moler mass. The
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 range of about 5 to 25 % by weight of the total weight of the
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EP 0 282 451 A2

Description

Destructurized starch and method for making same

The present invention refers to destructurized starch and to a method for making same,

It has recently become known that natural starch which is found in vegetable products and which contains a defined amount of water, can be treated at elevated temperature and in a closed vessel, thereby at elevated pressure, to form a met. The process is conveniently carried out in an injection moiding machine or extruder. The starch is fed through the hopper onto a rotating, reciprocating screw. The feed material moves along the screw towards the tip. During this process, its temperature is increased by means of external heaters around he outside of the barrel and by the shearing action of the screw. Starting in the feed zone and continuing in the compression zone, the particulate feed becomes gradually molten. It is then conveyed through the metering zone, where homogenization of the meter cours, to the end of the screw. The molten material at the tip can then be further treated by injection molding or extrusion or any other known technique to treat thermoplastic meters.

This treatment, which is described in the European Patent Application No. 84 300 940.8 (Publication No. 118 240) yields a destructurized starch. The reason for this being that the starch is heated above the melting and glass transition temperatures of its components so that they undergo an endothermic transition. As a consequence a melting and disordering of the molecular structure of the starch granule takes place, so that a destructurized starch is obtained.

Although such destructurized starch is useful in molding techniques and extrusion it has been found, that the molded parts show a relatively high incidence of surface defects and the processed materials generally have relatively lower extensibilities.

It has now been found, that the above mentioned disadvantages can be overcome by heating starch, having a water content in the range of 10 to 25%, in a closed volume to elevated temperatures in the presence of a chain scission catalyst until the mass average molar mass of the native starch is reduced by a factor within the range of 2 to 5000.

The starch material obtained according to the present invention also exhibits improved flow characteristics especially for the production of thin valled articles so that due to the improved processability, defective parts are minimized as well as necessary subsequent controls reduced. Also lower pressures can be used for processing e.g., in highciton modifiang processing as well as lower temperatures.

The present invention therefore refers to a method for producing destructurized starch, comprising a chemically non-modified starch material having a water content in the range of about 10 to 25 9b weight of the total weight of the composition, in the presence of a chain scission catalyst in a closed volume to an elevated temperature sufficient to form a thermoplastic met and continuing heating until the mass average molar mass of said starch material is reduced by a factor of 2 to 5000, compared with its original mass average molar mass.

The present invention also refers to destructurized starch obtained by this method.

The "starch composition" means the starch/water composition therein being included the optional additives as described later on.

The term destructurized starch has been explained above. Chemically non-modified starch as such includes for example native starch, was well as geletinized or cooked starch and includes generally carbohydrates of or natural, vegetable origin, composed mainly of amylose and/or amylopedin. It may be extracted from various plants, examples being potatoes, rice, staploca, corn, and oreals such as rye, oats and wheat.

Various values can be found in the literature for the mass average molar masses of different starches. The term 'its mass average molar mass is reduced by a factor of 2 to 5000 compared to its original mass average molar mass; is to be injectated as follows:

45 Starch as any polymeric material is made of macromolecules whose molar masses are not all the same. The molar mass of individual macromolecules cannot be measured but what can be measured is an average of the molar mass of all the molecules.

Among the absolute methods used for the determination of average molar mass, light scattering is most fevoured and has been applied to almost all conceivable types of macromolecules. Light scattering weighs the various molecules in a sample according to their masses and hence yields a mass average molar mass. Because of that, the method does not break down at very high molecular weights and this is a basic advantage over other methods. However, the light scattering measurement of mass average molar masses requires the dissolution of the polymers in a solvent.

In case of native starch for example, no solvent is available to achieve a convenient solution for the measurement. Consequently, he average molar mass or native starch must be determined by extrapplation from a plot correlating the light scattering mass swenage molar masses of lower molecular weight starches soluble in dimethysulphoxide (DMSO) as the solvent to their intrinsic visosolities in both DMSO/6 molar aqueous urea 9:1 and DMSO as solvents. The solvent DMSO/6 molar aqueous urea 9:1 can be used to dissolve native starch and to measure the intrinsic visosolity but cannot be used for flight scattering.

Therefore a linear plot log molar mass (as obtained from soluble lower molecular mass starches through light scattering) as a function of log $[\eta]$ (η - infinisci viscosity, as obtained from the Intrinsic viscosity measurement of the corresponding lower molar mass starches) permits to determine the average molar mass of native starch from its Intrinsic viscosity measurement by extrapolating the upper part of the plot to its

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intrinsic viscosity value.

When native starch is treated with the catalysts under the conditions claimed in the present invention, the starch macromolecules are split in a statistical manner and distinct from enzymatic degradation producing a low proportion of fragments of very low molar mass.

The splitting is due to the statistical breakdown of the α- 1,4 glycosidic bonds with the corresponding reduction of the average molar mass of the initial macromolecules.

Therefore the ratio of the average molar mass values of native starch and catalytically modified starch is a measure of the extent of the catalytically induced modification.

In the present case the weight average molar mass of potato starch was experimentally determined from intrinsic viscosity measurements to be about 6 × 10⁸. A reduction by a factor of 100 would therefore lead to a wei

weight average motar mass of about 6 × 10°.

The method of using light scattering to measure the mass average motar mass of starch is described in

Methods in Carbothydrate Chemistry, Vol IV, Starch, Academic Press, 1964, p. 191-202.
The use of the Intrinsic viscosity for molar mass determination of starch is described in Methods in Carbothydrate Chemistry, Vol IV, Starch, Academic Press, 1964, p. 179-188.

Carbohydrate Chemistry, Vol IV, Starch, Academic Press, 1964, p. 179-188.

Starch can also be characterized by its dextrose equivalent (DE), i.e. the granules of dextrose which have the same reducing sugar content as 100 grammes of the starch sample. The DE-values measured for the

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starch samples obtained according to the present invention were in the range of 40 to 700. The starch material has a water content in the range of about 10 to 25 % and better 10 to 20 % calculated to the weight of starch composition. Preferred is a final water content of 12 to 19 % and especially 14 to 18 % calculated to the weight of starch composition.

The starch material is heated to elevated temperatures sufficient to form a thermoplastic melt lac, temperatures higher than the glass transition and/or melting temperature of the formed material, temperatures are preferably in the range of 100 to 200°C, preferably 140 to 190°C. Most preferred are temperatures within the range of 160 to 185°C.

A thermoplastic met is a polymeric fluid having a shear rate dependent viscosity, as generally known. The starch melts according to this invention have similar magnitudes and dependences of viscosity on shear rate as conventional thermoplastic melts like o.g. those of polyethylene. This also means that viscosity decreases with increasing shear rate as well as with increasing temperature.

The starch material is heated preferably in a closed volume. A closed volume can be a closed vessel or the volume created by the sealing action of the unmoiten feed material as happens in the screw of injection moiding or extrusion equipment. In this sense the screw-barrel of an injection moiding machine or an extruder is to be understood as being a closed vessel, Fessures created in a closed vessel correspond to the vapour pressure of water at the used temperature but of course pressure may be applied as this is normally done in a screw-barrel. The preferred applied pressures to be used are in the range of the pressures which are applied in extrusion or injection moiding processes and known per se, i.e. from zero to 150 × 10° N/m² a.

The melt of destructurized starch according to this invention is e.g. injected under the normal range of pressures used in injection molding namely for thicker walled articles in the range from $300 \times 10^5 \text{ N/m}^2$ to $3.000 \times 10^5 \text{ N/m}^2$, preferably $700 \times 10^5 \times 2000 \times 10^5 \text{ N/m}^2$.

A chain scission catalyst is a catalyst which acts by splitting mainly the α - 1,4 glycosidic bonds in the starch macromolecule in a statistical manner regular manner.

Useful catalysts are acids and bases including Lewis acids and bases or any reagent which can catalyse the splitting of the α - 1,4 glycosidic bond in a statistical manner.

Suitable acids are inorganic acids such as hydrochloric acid, suffuric acid, nitric acid, phosphoric acid, are horizonate acid, hydrofluoric acid, hydrofluoric mous acid, hypochrousus acid, hypochrousus acid, periodic acid, aci

Sultable are further organic acids such as formic acid, acetic acid, propionic acid, n-butyric acid, halogenacette acid, lactic acid, sybcoile acid, oxalic acid, matonic acid, citric acid, tartaric acid, itaconic acid, succinic acid, glucoric acid, guitaric acid, acid, pimelic acid, organo-sutrionic acid, berarcio acid, ethylenediamine tetracetic acid (EDTA), acuste amine acides such as guitaric acid, aspartic acid, or a mixture of these compounds as well as partial salts of the polyacids mentioned, e.g. discolum EDTA.

Preferred are hydrochloric acid, sulfuric acid, disodium-EDTA, citric acids or a mixture of these compounds, Sultable bases are inorganic bases such as sodium hydroxide, potassium hydroxide, calcium hydroxide, acicium hydroxide, magnesium hydroxide, ammonia, aluminium hydroxide, hydrazine, or a mixture of these compounds.

Suitable are further organic bases and Lewis bases such as metriylamine, etrylamine, propylamine, butylamine, and homologues, anline, benzylamine, ethanediamine, propanediamine, butanediamine hexametrlylenediamine, morpholine, pryidine, piperidine, basic amino acids such as arginine, asparagine, glutamine, histoline, trytophan or a mixture of these compounds.

Preferred are ammonia, sodium hydroxide, potassium hydroxide, calcium hydroxide, magnesium hydroxide, tryptophan or a mixture of these compounds.

The acids mentioned above are the preferred catalysts. The reduction of the mass average molar mass of the original (chemically non-modified) starch used is by the factor 2 to 5000, preferably by a factor of 4 to 1000 and most preferably by a factor of 5 to 500.

The usual heating time is about 2 - 10 minutes and preferably 4 - 8 minutes, but this time Indication is not critical and can be varied.

The concentration of catalysts is in the range of 10^{-8} to 10^{-2} mole of catalyst per mole of anhydro-glucose unit (AGU), preferably between 0.1×10^{-3} to 5×10^{-3} mole/AGu.

The starch material of the present invention may contain or may be mixed with additives such as extenders, lubricants, plasticers and/or coloring agents. Such materials are named in the European Patent Application

No. 84 300 940.8 (Publ. No. 118 240) and are incorporated herewith by reference.
In this sense, these additives may be added before, during or after heating the starch to form the melt. After

heating would mean e.g. when the melt is cooled down before further processing. It mainly depends on the intended use of the destructurized starch.

Such additives are extenders of different kinds, e.g. gelatin, vegetable proteins such as sunflower protein, soybean proteins, cotton seed proteins, peanut proteins, rape seed proteins, blood proteins, egg proteins, acrylated proteins; water-soluble polysaccharides such as:

alginates, carrageenans, guar gum, agar-agar, gum arabic and related gums (gum ghatti, gum karaya, gum tragacauth) pectin; water-soluble derivatives of cellulose:

20 ali/cielluloses hydroxyalkyloelluloses and hydroxyalkyloelluloses, such as: methyloellulose, hydroxyar-methyloellulose, hydroxyethyloellulose, hydroxyethyloellulose, hydroxyethyloellulose, hydroxyethyloellulose, hydroxyethyloellulose, hydroxyethyloellulose, hydroxyethyloellulose, bydroxyalkyloelluloseseters such as: cellulosesectylphitalate (CAP), hydroxyproythertyloellulose, HydroXy), activoxyalkyloellulose, activoxyalkyloellul

alkylcelluloses, carboxyalkylcelluloseesters such as:

25 carboxymethylceliulose and their alkalimetal salts; water-soluble synthetic polymers such as; polyacrylic acids and polyacrylic acids esters, polymethacrylic acids and polymethacrylic acid seters, polyminylacetates, polyminylacetates,

Such extenders may optionally be added in any desired amount preferably up to and including 50 %, preferably within the range of 3 % to 10 % based on the weight of all components.

The choice of catalyst and extender should be done in such a way that both are compatible i.e. that the catalyst cannot be harmful to the extender and also retains its own activity. This principle is of course applied also with respect to the additives mentioned further or the course of the co

Further additives are inorganic fillers, such as the oxides of magnesium, aluminum, silicon, titanium, etc. preferably in a concentration in the range of about 0.02 to 3 % by weight preferably 0.02 to 1 % based on the weight of all components.

Further examples of additives are plasticizers which include polyalkylene oxides, such as polyethylene glycols, polypropylene glycols, organic plasticizers with low molecular weights, such as glycerol, glycerol monoacetate, diacetate or triacetate; propylene glycol, sorbidt, sodium diethylsulfosuccinate, triethyl citrate, triiutyl citrate, etc., added in concentrations ranging from 0.5 to 15 %, preferably ranging from 0.5 to 5 % based on the weight of all the components.

Examples of coloring agents include known aco dyes, organic or inorganic pigments, or coloring agents of natural origin, inorganic pigments are preferred, such as the oxides of iron or trianium, these oxides, known per se, being added in concentrations ranging from 0,001 to 10 %, preferably 0.5 to 3 %, based on the weight of all the components.

The sum of the plasticizer and water contents should preferably not exceed 25 %, and should most preferably not exceed 20 %, based on the weight of all the components.

There may further be added compounds lubricants to improve the flow properties of the starch material such as animal or vegetable fats, preferably in their hydrogenated form, especially those which are solid at room temperature. These fats have preferably a melting point of 50°C or higher. Preferred are Triglycerides Wiffin C12 - C14 - C14 - C16 - 181 (21-181) and C15.

These fats can be added alone without adding extenders or plasticizers.

These fats can advantageously be added alone or together with mono- and/or diglycerides or phosphatides, especially lecithin. The mono- and diglycerides are preferably derived from the types of fats described above, i.e. with Ct₂ -, Ct₄ -, Ct₉ -, and Ct₉ - fatty acids.

The total amounts used of the fats mono-, diglycerides and/or lecithins are up to 5 % and preferably within the range of about 0.5 to 2 % by weight of the total composition.

It is further recommended to add silicon dioxide or titanium dioxide in a concentration of about 0.02 to 1 % by weight of the total composition. These compounds act as texturizing agent.

by weight of the present invention, the materials treated as described herein above form on heating and in a closed vessel a melt with thermoplastic properties, i.e. under controlled water-content and pressure conditions, Such a melt can be used in various techniques just like thermoplastic materials. These techniques

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include injection molding, blow molding, extrusion and coextrusion (rod, pipe and film extrusion, compression molding, accuum forming, to produce known artibles as produced with these techniques include bottles, sheets, films, packaging materials, pipes, rods, laminates, sacks, bags, pharmaceutical capsules.

The following examples further explain the invention.

Example

Natural potates stams, Type AVEBE, a lubricant/release agent (typdrogenated fat) and a melt flow accelerator (ecithin), a texturizing agent (TiQ₂) and hydrochoric acid (1 moiar) are mixed together in the relative proportions in a high speed powder mixer for 10 minutes so that a composition consisting of 92.9 parts of natural potate stame, 0.83 parts of the hydrogenated triglycentide containing the fatty acids C_{18} : C_{18} : C_{18} : C_{18} is a ratio of 55:314 weight percent, 0.415 parts techtin, 0.415 parts techting technique techting technique technique

This powder is fed into the hopper and fed to the screw barrel having a temperature of 170 - 180°C and an applied pressure of 55 - 75 bar and kept at these conditions for 7 minutes. It was the ninjection molding tool for capsule body and cap parts whose mold wall temperature was kept at 40°C using an injection molding pressure of 1000 bar. After cooling and ejecting from the mold a dimensionally stable capsule body and cap part is obtained which can easily be processed on a filling machine.

The analysis of the molar mass was done by determination of the viscosity and correlation with the light scattering measurements. The reduction of the molar mass was a factor of 4,7 and gives an improvement of a capsule quality down to 1 % defects compared to 9 % obtained from the above composition without having added HCl to 1.

Example 2

Natural potato starch Type SUD with the same additives as in Example 1 and 0,5 × 10⁻³ mole hydrochloric acid/AQU is processed in the same manner as in Example 1 (Temp: 170°C, 50 bar pressure, 7 minutes). The analysis of the molar mass was done by determination of the viscosity and correlation with the light scattering masurements. The reduction of the molar mass was a factor of 2,5 and gave an improvement of capsule quality down to 0.7 % of defects compared with 5 % of defects obtained with the same starch composition without having added the each. Analogous results are obtained with fice starch.

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Table 1 Examples 3 to 16)

In the following table 1 there are given Examples where potato starch has been mixed with the same additives as described in the Examples 1 and 2 but processed undow revious add contents, temperatures and residence times, obtaining different values for the reduction factors of the mass average molar masses of the starches used. The applied pressures were as described in Example 1. Considerable improvements in he numbers of defects were consistently obtained. The experiments also showed, that it is possible to reduce the numbers of defects were consistently obtained. The experiments also showed, that it is possible to reduce the names average molar mass by healing starch at elsevated temperatures and pressures without the addition of a chain solision catalyst. However, this reduction alone does not give the consistent reduction in defects. Surprisingly, the presence of the chain solision catalyst in combination with the reduction in the mass average molar mass to values within the limits of 2 to 5000 as described above are necessary to obtain the results of the present invention.

It is well known that different samples of natural starch materials show variations in their structures and properties. The consistent reductions in defects surprisingly occur in spite of such variation differences in the structures and properties of the granules of the various native potato starches used.

Table 2 shows further compositions that are treated analogously as described in the Examples 1 or 2.

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Number of defects in	5.0	3.7.	2.7	68 3.5	5.0	9.3	3.2	
Reduction factor MM native starch MM product	184 475	162 330	211 260	24.5	52 259	69.7	72.3	
Residence time in the screw barrel (nozzle) sec.	475 475	665 590	665 590	965 590	500 450	665 590	965 590	
Temp. in the screw barrel	170 165	160 155	160 150	160 155	170 165	160 155	160 155	
Catalyst con- centration mol/AGJ	0.5 × 10 ⁻³	0.5 x 10 ⁻³	0.6 x 10 ⁻³	0.7×10^{-3}	0.8 x 10 ⁻³	0.8 x 10 ⁻³	0.85 x 10 ⁻³	
Scission catalyst	(a) none (b) HCl	(a) none (b) H ₂ SO ₄	(a) none (b) H ₂ SO ₄	(a) none (b) H ₂ SO ₄	(a) none (b) HCl	(a) none (b) H ₂ SO ₄	(a) none (b) H ₂ SO ₄	
Type of starch: potato	Südstärke Batch 1	Roquette Batch 2	Roquette Batch 7	Roquette Batch 5	Avebe Batch 2	Roquette Batch 6	Roquette Batch 9	
Example No.	e	4	ري د	9	7	ω	o,	

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No. STARCH	COMPONENT b)		COMPONENT c)		COMPONENT d)		WATER	OTHERS	
Type 8	Type	*	Type	æ	Type	مو	οp	Type	esp.
ĺ				,	C Fill	2	9 %		
Potato 75.00	hydrated vege- table fat	1.25	Lectthin	ç	2021	0.63	2.3		
Potato 80.00	•	1.0		6.0	•	0.1	18.0	ı	
Potato 88.6	:	9.0		0.5	sto ₂	0.1	10.0	ı	ı
Wheat 78.0			Monoglycerid Cle	5.0		3.04	19.0	dyestuff	96.0
Wheat 79.0		2.05	•	,	710, S10, 1:1	0.95	18.0	,	1
Wheat 85.0	hydrated animal fat	2.80	ı	,	110 ₂	0.20	12.0	1	ı
Maize 75.0		4.80	Monoglycerid C ₁₈	0.2		0.15	17.0	pre-cocked starch	2.85
Maize 78.0	:	3.0		1.0		0.02	14.0		3.95
		1		5.0	StO2	0.7	17.5	dyestuff	5.0
Rice 80.25	‡	6.0	Diglyceride	9.0		0.25	0.51	Glyœrdn	m
Taploca 75.2		6.0		9.0		0.3	21.0	HAPCP	2
		2.0		2.0		0.04	17.0	PEG 1000	S
	r.	1.5		1.0	${\rm THO}_2$	0.25	18.0	PEG 4000	4
Potato 75.25	5 vegetable fat	1.5	Lecithin	1.0		0.25	18.0	Sorbitol	4
		1.5		1.2		0.25	17.0	Polyacrylic 20.05 acid	20.05

TABLE

Claims

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- 1. A method for producing destructurized starch, comprising heating a chemically non-modified starch material having a water content in the range of about 10 to 25 % by weight of the total weight of the composition, in the presence of a chain scission catalyst in a closed volume to an elevated temperature sufficient to form a thermoplastic meth and continuing heating until the mass average molar mass of said starch material is reduced by a factor of 21 to 500.0 compared with its oriolinal mass average molar mass.
- 2. A method according to claim 1, wherein said chemically non-modified starch is at least one member of a class selected from native starch, gelatinized or cooked starch or generally carbohydrates of natural, vegetable ordinic, composed mainly of amylose and/or amylosectin.
 - 3. A method according to the claims 1 or 2, wherein the starch material has a water content in the range of about 10 to 25 % and better 10 to 20 % calculated to the weight of the starch composition preferably a water content of 12 to 19 % and especially 14 to 18 % calculated to the weight of starch composition.
 - 4. A method according to anyone of the claims 1 to 3, wherein the starch material is heated to an elevated temperature, preferably in the range of 100 to 200°C, preferably 140 to 190°C, and especially to
- temperatures within the range of 160 to 185° C.

 5. A method according to anyone of the claims 1 to 4, wherein pressure is applied in the range of from zero to 150 × 10° N/m² and most particularly from zero to 50 ×
- 10⁸ N/m².

 6. A method according to any one of the claims 1 to 5, wherein a chain scission catalyst is added which acts by splitting the -1,4 glycosidic bonds in the starch macromolecule in a statistical or regular manner,
- preferably an acid or a base, including Lewis acids.

 7. A method according to claim 6, wherein the chain scission catalyst contains an inorganic acid, a
- Lewis acid, an organic acid or a mixture of such compounds.

 8. A method according to claim 6, wherein the chain solssion catalyst contains an inorganic base, an
- organic base, a Lewis base or a mixture of such compounds.

 9. A method according to claim 6, wherein said chain scission catalyst is selected from hydrochloric acid, suffuric acid, EDTA, citric acids or a mixture of these compounds.
 - 10. A method according to claim 6, wherein said chain scission catalyst is added in a concentration in the range of 10 $^{-2}$ to 10 $^{-2}$ mole of catalyst per mole of anhydro-glucose unit (AGU), preferably between 0.1 \times 10 $^{-3}$ to 5 \times 10 $^{-3}$ mole/AGU.
 - 11. A starch material according to anyone of the claims 1 to 10, characterized in that the reduction of the mass average molar mass of the starch used is by the factor 2 to 5000, preferably by a factor 4 to 1000 and
 - most preferably by a factor 5 to 300.

 12. A method according to anyone of the claims 1 to 12, characterized in that one or more additive(s) is/are added before, during or after heating of the starch material to form a thermoplastic melt.
 - 13. A method according to claim 12, characterized in that there is added at least one extender or a mixture of extenders within the range of up to 50 %, preferably within the range of 3 % to 10 %, based on the weight of all components.
 - 14. A method according to the claims 12 or 13, characterized in that there is added at least one inorganic filler or a mixture of such fillers in a concentration of about 0.02 to 3 %, preferably 0.02 to 1 %, by weight of all components
 - 15. A method according to anyone of the claims 12 to 14, wherein a plasticizer within the range of about 0.5 to 15 %, preferably 0.5 to 5 % by weight of all components.
 - 16. A method according to anyone of the claims 12 to 15, wherein a coloring agent is added in a concentration of about 0.001 to 10 %, preferably 0.5 to 3 % by weight of all components.
- 50 17. A method according to anyone of the claims 12 to 16, wherein a plasticizer is added and the sum of the plasticizer and water contents does not exceed 25 %, and preferably does not exceed 20 %, by weight of all components.
 - 18. A method according to anyone of the claims 12 to 17, wherein a lubricant is added to improve the flow-properties optionally together with a monoand/or diglyceride and/or a phosphatide in an amount of up to 5 %, preferably within the range of 0.5 to 2 % by weight of the total composition.
 - 19. A method according to anyone of the claims 12 to 18, wherein there has been added silicon dloxide or titanium dioxide has been added in a concentration of about 0.02 to 1 % by weight of the total composition.
 - 20. A destructurized starch obtained according to anyone of the claims 1 to 19.
 - 21. The process of shaping destructurized starch according to anyone of the claims 1 to 19, under controlled water content and pressure conditions as a thermoplastic melt where insaid shaping process is at least one member selected from the class consisting of injection molding, blow molding, extrusion and coextrusion, compression molding or vacuum forming.
 - Shaped articles produced from a composition obtained by the method as claimed in anyone of the claims 1 to 19.

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23. The articles of claim 21, wherein said articles have been shaped as bottles, sheets, films, packaging materials, pipes, rods, laminates, sacks, bags or pharmaceutical capsules.